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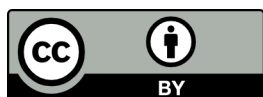
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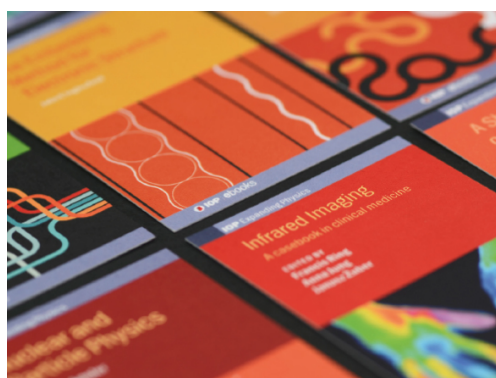
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Determination of the preferred reaction pathway of acetophenone on Si(001) using photoelectron diffraction

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Abstract

The adsorption configurations of a technologically relevant model organic adsorbate on the silicon (001) surface were studied using energy scanned x-ray photoelectron diffraction (PhD). Previous work has established the existence of an interesting vertically-aligned ('flagpole') configuration, where the acetophenone attaches to Si(001) via the acetyl group carbon and oxygen atoms. Density functional theory calculations have predicted two energetically similar variants of this structure, where the phenyl ring is orientated parallel or perpendicular to the rows of silicon dimers on this reconstructed surface. However, previously published experimental measurements, including scanning tunnelling microscopy, x-ray photoelectron spectroscopy, and near-edge x-ray absorption fine structure investigations were unable to distinguish between these two configurations. Here, we apply the unique experimental capabilities of the PhD technique to this system and demonstrate that the dominant adsorption configuration has the phenyl ring parallel to the dimer rows (the end-bridge structure). This information in turn facilitates the determination of the dominant reaction pathway for acetophenone on Si(001), which has remained elusive until now. Information about subtle preferences in reaction pathways that affect the alignment and orientation of organic adsorbates such as acetophenone on technologically-relevant semiconductor surfaces such as

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Si(001) is critical for the fabrication of future atomically-precise atomic and molecular-scale electronic devices utilising the organic-silicon interface, and this work demonstrates the unique and complementary capabilities of PhD for providing this information.

Keywords: structure determination, organic-silicon interface, Si(001), energy scanned photoelectron diffraction, density functional theory

 Supplementary material for this article is available [online](#)

(Some figures may appear in colour only in the online journal)

1. Introduction

As fundamental length scale limits are being reached in the utilisation of silicon in the semiconductor industry, the pursuit of novel electronic devices has become a necessity. With a view to shrinking the size of electronic components even more, the organic-silicon interface has drawn significant attention to the development of miniature devices that function on the molecular level [1–6]. However, incorporating molecular components into silicon-based technology presents many challenges. For example, molecules that only bind weakly to the substrate result in structures that are unstable or the silicon surface being incompletely passivated, with the resulting dangling bonds leading to deleterious electronic effects [7]; alternatively, poor electronic overlap between the molecular orbitals and the band structure of the substrate can necessitate high electric fields to align the two, resulting in unwanted dynamic changes to the molecular structure [1, 8]. Thus, developing methods to robustly attach molecules in precise adsorption configurations with appropriate electronic coupling to the substrate and that are robust under applied electric fields remains a significant challenge.

The ambiphilic nature of the Si(001)-(2 × 1) reconstruction involves buckled dimers with a charge transfer of about 0.36 electrons from the down-buckled to the up-buckled dimer atom [9–11]. The electron-poor down-buckled atom presents a strongly reactive site for nucleophiles such as methanol [12] and phosphines [13], while select electrophiles such as borane have been shown to react with the electron-rich up-buckled atom [14]. Molecules containing carbonyl bonds constitute a particularly interesting class of adsorbate species because, during a sequence of reaction steps, they behave both as nucleophiles through the lone pair of the carbonyl oxygen, and electrophiles through the electron-deficient carbonyl carbon [15]. Moreover, acetone and acetophenone have been shown to form a strong O–Si bond with the silicon substrate, forming structures that are stable under thermal annealing and within a high local electric field, e.g. between the surface and the tip of a scanning tunnelling microscope (STM) [16, 17].

Our prior work using near edge x-ray absorption fine structures (NEXAFS) [18], density functional theory (DFT), and STM [16, 17] to study the adsorption of acetophenone on Si(100) demonstrated that, after room temperature deposition, the phenyl ring of the molecule is found to align mostly parallel

to the surface plane with two different orientations (figures 1(a) and (b)). These flat lying molecules were assigned to an allyl structure, where the aromaticity of the phenyl ring is broken by coordination of the two carbon atoms of the phenyl ring to two silicon atoms of the surface. Two distinct orientations of this allyl species were assigned to two different anchoring geometries of the terminal oxygen and carbon atoms of the acetyl group to the surface silicon dimers. Specifically, the terminal atoms bridging two atoms from different silicon dimers (end bridge allyl, EB-allyl, see figure 1(a)), or anchored to two atoms from the same silicon dimer (dimer bridge allyl, DB-allyl, see figure 1(b)). We found that the initial allyl adsorption structure could be manipulated into a structure with greater thermodynamic stability: after mild annealing, or via application of a voltage/current pulse from the STM tip, the adsorbate changes configuration such that the phenyl ring is mostly perpendicular to the surface plane, referred to as the flagpole structure. This transition is also deduced from the carbon 1s x-ray photoelectron spectroscopy (XPS) data, where a new peak appears at lower binding energy due to hydrogen dissociation from the methyl group [18]. This flagpole structure is both thermodynamically stable, and exhibits strong electronic coupling to the silicon substrate [16, 18], making this adsorbate desirable for device applications, or as an anchoring group for further functionalisation.

As with the allyl structure, two orientations of the flagpole structure with respect to the surface silicon dimers are possible: bridging across two Si dimers (end bridge flagpole, EB-flagpole, see figure 1(c)), or bridging a single Si dimer (dimer bridge flagpole, DB-flagpole, see figure 1(d)). Knowledge of the orientation of this flagpole structure is important in the context of molecular device fabrication for two reasons: (1) it provides a means to determine experimentally the preferred reaction pathway on the silicon surface, which is important not only for this adsorbate, but the class of adsorbates that anchor to silicon via an acetyl group; and (2) the orientation of the adsorbate with respect to the surface dimers can be a critical factor in the further functionalisation of the surface [19]. While STM is capable of differentiating between the two orientations of the allyl features (figures 1(a) and (b)), the large aspect ratio of the two flagpole structures (figures 1(c) and (d)) makes differentiation between these two features impossible using STM. NEXAFS is similarly incapable of determining

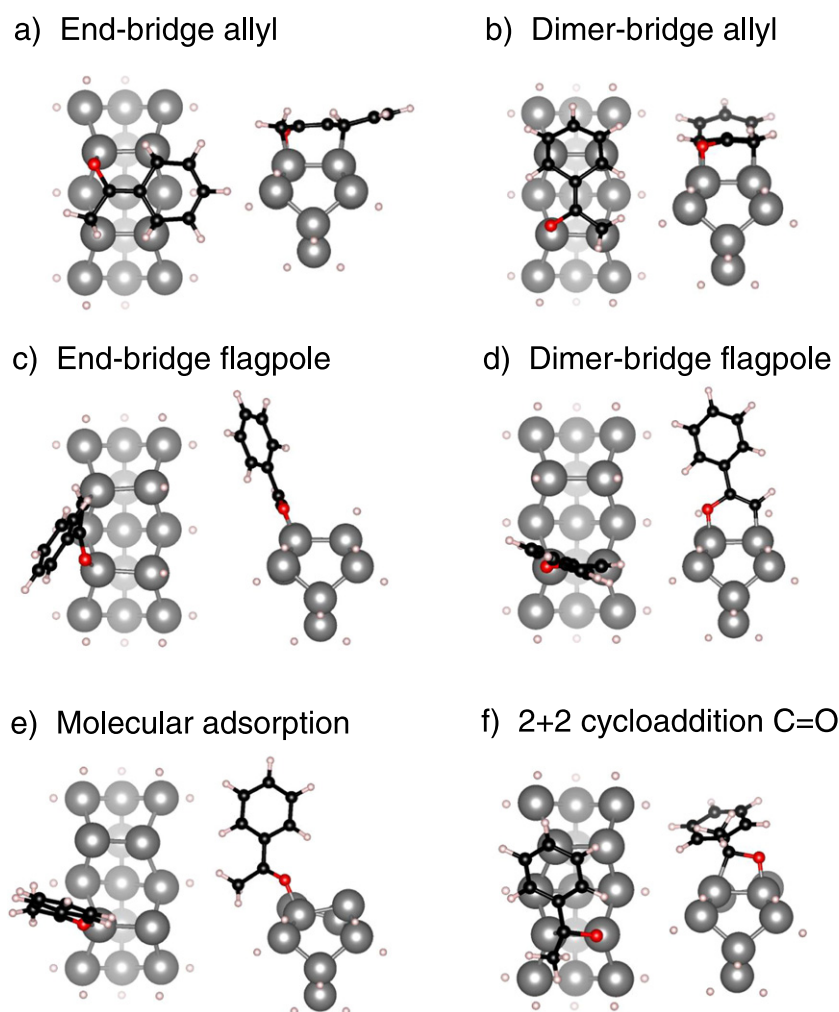


Figure 1. Top and side views of the structures predicted by DFT: (a) EB-allyl, (b) DB-allyl, (c) EB-flagpole, (d) DB-flagpole, (e) molecular adsorption, (f) 2 + 2 cycloaddition C=O. The black atoms are C, white H, red O and grey Si. In the plan views the Si atoms below the dimers are shaded in a lighter colour as a function of depth.

the azimuthal orientation of the molecule and thus differentiating the two models [20] due to the four fold symmetry of the surface, nor can these structures be differentiated using XPS.

For completeness, in this work we also consider two additional structures that were not part of earlier DFT calculations. These are the molecularly adsorbed configuration, shown in figure 1(e) and a C=O 2 + 2 cycloaddition structure shown in figure 1(f). These structures were modelled, since we considered it plausible that they might produce a fit to the photoelectron diffraction (PhD) data; however, as described below, this did not turn out to be the case and these structures were ruled out.

Energy-scanned x-ray PhD is arguably the only technique with the potential for experimentally determining the structural properties of the acetophenone flagpole structure on Si(001). This technique exploits the coherent interference of the directly-emitted photoelectron component, and components of the same photoelectron wavefield elastically backscattered by neighbouring atoms [21]. As the photon energy,

and therefore the photoelectron kinetic energy/photoelectron wavelength, is varied, different scattering pathways come into and out of phase, yielding modulations in the photoemission intensity. These modulations contain information about the relative distances and orientation between the emitter and the scatterers, and allow us to obtain precise structural information without requiring long range order.

In this work, we explore the powerful and unique capabilities of PhD for determining detailed structural information for a model adsorbate (acetophenone) adsorbed on the technologically ubiquitous silicon (001) surface. In particular, we measure silicon (001) surfaces exposed to acetophenone before and after being subjected to mild thermal annealing. Using this method, we determine the preferred orientation of the acetophenone flagpole structure, thereby elucidating the preferred kinetic pathway of the acetophenone adsorbate among those that have been theoretically predicted [16, 17]. We also discuss the limitations of this technique for the precise determination of structural parameters of organic adsorbates on silicon.

2. Experimental and computational details

2.1. Experimental details

The experiment was performed at the PEARL beamline [22] of the Swiss Light Source. The end-station of the PEARL beam line consists of an analysis chamber, a low-temperature STM chamber and a preparation chamber. All chambers have a base pressure $\sim 3 \times 10^{-10}$ mbar. A clean Si(001) (As-doped, 0.0015–0.0400 Ω cm, Virginia Semiconductor) surface was prepared by repeated resistive annealing of the silicon crystal with 45 W (6 A/7.5 V) for 20 s with 1 min ramp down from 2 A, with the chamber pressure staying below $\sim 5 \times 10^{-9}$ mbar. The crystalline quality of the surface was judged by STM measurements; the cleanliness of the surface by synchrotron XPS (SXPS). Acetophenone was exposed to the sample through a leak valve at a partial pressure of 1×10^{-8} mbar for 300 s (~ 3 L, where 1 L is 1×10^{-6} mbar s). Prior to exposure the gas line between the acetophenone container and the leak valve was baked and then conditioned under a flow of acetophenone. Prior to annealing the acetophenone layer, the sample was re-prepared with a new acetophenone layer in order to avoid effects from potential beam damage. To anneal the acetophenone layer, the Si sample was again resistively annealed but at a lower power of 1.11 W (3.7 V/0.3 A) for 10 min. Oxygen 1s, carbon 1s and silicon 2p SXPS measurements were taken of both preparations [shown in the ESI, figure S1 (<https://stacks.iop.org/JPCM/33/214002/mmedia>)], and conform well to our previously published results [18].

The XPS and PhD measurements were acquired from a Scienta EW4000 hemispherical electron energy analyser. The analyser was mounted at an angle of $\sim 60^\circ$ with respect to the incident beam in the plane of the photon polarisation (linear horizontal). The slits of the detector were orientated perpendicular to polarisation of the incident light and, as such, for the PhD measurements it was considered as a point detector utilising only the inner $\pm 5^\circ$ of the analyser's acceptable angle. The end-station is equipped with a six-axis manipulator, allowing translations in x , y and z and polar, azimuthal and tilt rotations.

Oxygen 1s PhD modulation spectra were recorded for both preparations in the photon energy range of ~ 630 – 890 eV, with an energy step of 4 eV. The data were acquired along the [110] and [100] crystallographic directions in the polar emission angle range of 0° – 60° . The resulting energy distribution curves were processed using our standard methodology [21, 23], whose details can be found in the ESI. Carbon 1s PhD modulation spectra were also acquired (see figure S2 in the ESI), but no significant modulations were observed, thus fitting of this data was not pursued and the quantitative analyses were based solely on the O 1s PhD modulations. A full list of the fitting parameters that were allowed to vary in the calculations is detailed in the ESI. The three fitting parameters that were found to strongly affect the fitting were those defining the bond vector between the oxygen atom and the silicon atom it was bound to. These positions were defined in spherical coordinates: bond length (R), tilt of the bond away from the surface normal (θ) and the rotation of the bond away from the [110]

direction (ϕ). The other fitting parameters are not discussed as, individually, they had negligible effect on the quality of the fit.

To compare the agreement between the resulting PhD and the DFT structures a fitness parameter, ΔF , was introduced:

$$\Delta F = \sqrt{\sum_i F_i^2}, \quad (1)$$

$$F_i = \frac{X_{(i)}^{\text{PhD}} - X_{(i)}^{\text{DFT}}}{\sigma_{(i)}}. \quad (2)$$

Here $X_{(i)}^{\text{PhD}}$ and $X_{(i)}^{\text{DFT}}$ are the values of structural parameter i of the PhD and DFT structures, respectively; σ_i is the associated uncertainty in that fitting parameter from the PhD structure with one of the best found R -factor, R_{fac} . The three structural parameters used for this calculation were R , θ and ϕ , with associated uncertainties of 0.05 Å, 4° and 12° , respectively. Any structure with a value of ΔF greater than 2 is considered significantly different from the DFT model.

2.2. Computation details

DFT calculations were performed using the Gaussian 09 suite of programs [24]. The surface was modelled by a $\text{Si}_{15}\text{H}_{16}$ cluster that represented two silicon dimers within the same row. Calculations were performed using the B3LYP method [25, 26] with the 6-311 + $G(d, p)$ basis set paired with Grimme's D3 dispersion corrections [27] and original damping function. The coordinates of atoms representing third through fifth rows of silicon atoms in the model clusters were fixed at bulk positions to avoid unrealistic distortion. After the $\text{Si}_{15}\text{H}_{16}$ cluster was optimized, the appropriate adsorbate was placed on the sites representing the surface and the structure was reoptimized at the same level of theory and with the same geometrical constraints. Adsorption energies were calculated from total energy calculations according to the formula:

$$E_{\text{adsorption}} = E_{\text{adsorbate+substrate}} - (E_{\text{substrate}} + E_{\text{gas phase molecule}}). \quad (3)$$

In the case of the two allyl structures, adsorption of the molecule to the surface is dissociative, resulting in the formation of a surface silicon hemihydride dimer. In these two cases, the adsorption energy was calculated using:

$$E_{\text{adsorption}} = E_{\text{adsorbate+substrate}} + E_{\text{hemihydride}} - (2E_{\text{substrate}} + E_{\text{gas phase molecule}}). \quad (4)$$

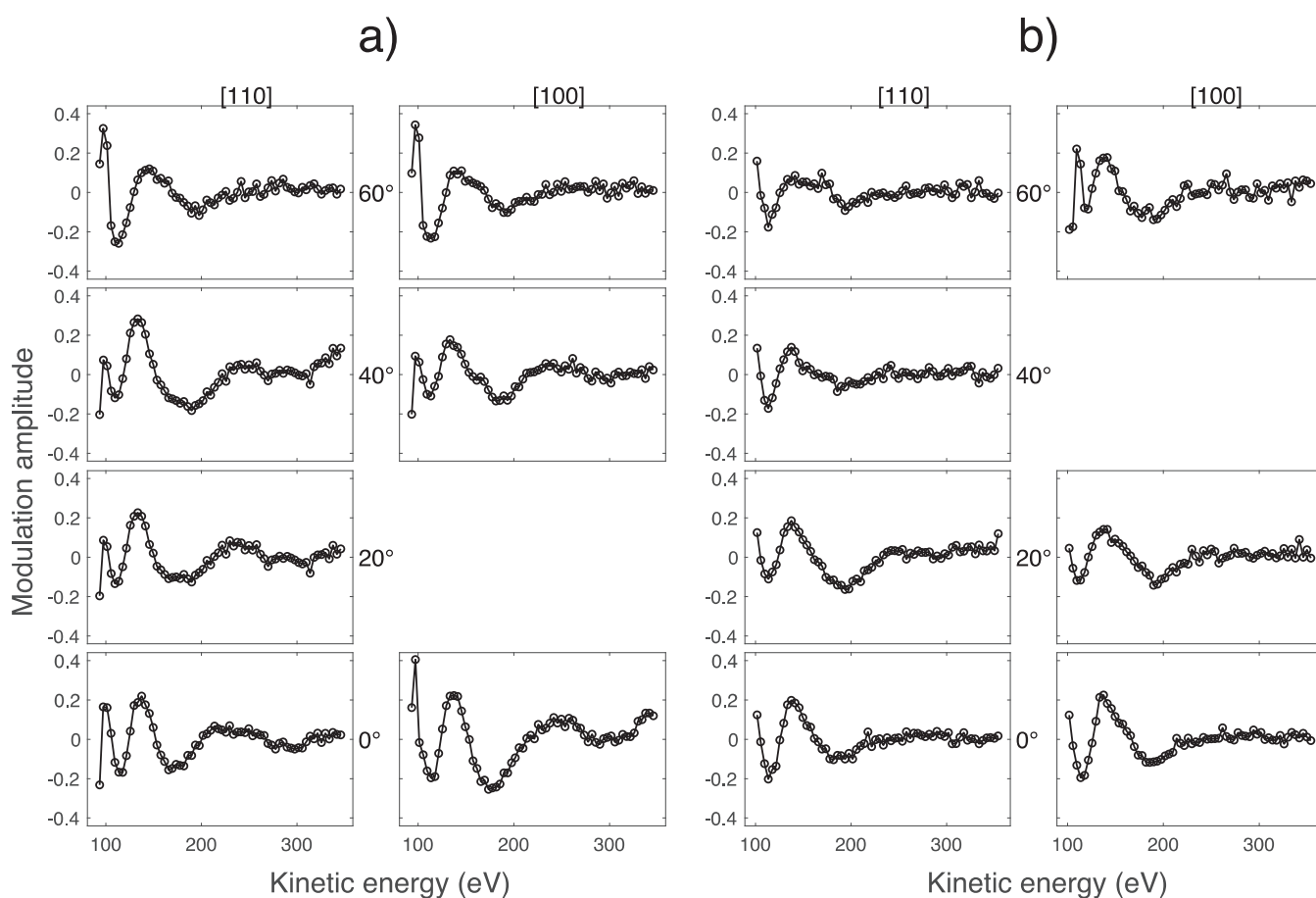
3. Results

3.1. DFT modelling

Six unique structures were modelled in this study. Three of the models are those presented in our prior work [16, 18]: EB-allyl (figure 1(a)), DB-allyl (figure 1(b)) and DB-flagpole (figure 1(d)). We also present calculations of the EB-flagpole (figure 1(c)), in addition to two wholly novel structures: a molecularly adsorbed structure (figure 1(e)) and a 2 + 2 cycloaddition structure (figure 1(f)). These two novel

Table 1. Calculated adsorption energies for the structures shown in figure 1.

	Molecular	2 + 2 C=O	EB-allyl	DB-allyl	EB-flagpole	DB-flagpole
Energy (eV)	−1.12	−2.02	−3.39	−3.27	−3.76	−4.15

**Figure 2.** Oxygen 1s modulations amplitudes for (a) the room-temperature and (b) annealed preparation. The azimuth is shown in each column and polar emission angles with respect to normal emission in each row.

structures were found to be significantly less energetically favourable than the allyl and flagpole structures (see table 1). Notably the energy minimised molecular adsorption structure is orientated almost perpendicular to the surface, tilted 83° away from the surface plane, which would exclude it from being the primary structure present after deposition at room temperature [16, 17]. Furthermore, the methyl group is not deprotonated in this structure, hence it would not account for the appearance of the lower binding energy peak in the C 1s XPS spectrum after annealing. The 2 + 2 cycloaddition structure has the carbon and oxygen atom of the carbonyl bound to two separate silicon atoms. This configuration has the phenyl ring almost parallel to the substrate, tilted 26° away from the surface plane, which would be consistent with the room temperature deposition phases; however, it is unclear how such a configuration would result in the double lobed structure observed in STM, and is over 1 eV less favourable than either allyl structure. Regardless of this apparent predisposition towards the allyl and flagpole structures, the

molecular adsorption and the 2 + 2 cycloaddition structures were also considered in the PhD analysis.

The two allyl structures (end bridge and dimer bridge; EB and DB) have the terminal carbon atom of the acetyl group singly deprotonated; while the two flagpole structures (also EB and DB) have the terminal carbon atom doubly deprotonated. Consistent with prior work, we find here that the flagpole structures are overall more stable than the allyl structures, and the DB-flagpole model is predicted to be the most energetically favourable, with an adsorption energy 0.39 eV lower than that of the EB-flagpole (table 1). The DB- and EB-allyl have a far more comparable adsorption energy, only differing by 0.12 eV. The EB- and DB-allyl structures have the phenyl ring orientated more parallel to the substrate, tilted 15° and 33° away from the surface plane, respectively. The EB- and DB-flagpole structures, instead, have the phenyl ring almost perpendicular to the substrate, tilted 65° and 77° away from the surface plane. Note that in our prior NEXAFS study [18], we showed that at

Table 2. Best fitting PhD structural models for single site occupation of a given DFT model for the PhD data taken after room-temperature deposition. The structural parameters of the PhD fit are given, the corresponding parameters from the DFT calculations are shown in brackets.

	R (Å)	ϕ (°)	θ (°)	R -factor	ΔF
EB-allyl	1.65 ± 0.06 (1.75)	55 ± 15 (66)	19 ± 4 (20)	0.45	2.2
DB-allyl	1.65 ± 0.05 (1.74)	83 ± 18 (83)	15 ± 5 (14)	0.46	1.9
EB-flagpole	1.68 ± 0.06 (1.71)	39 ± 19 (39)	23 ± 4 (26)	0.44	1.0
DB-flagpole	1.63 ± 0.05 (1.72)	16^a (10)	12 ± 7 (5)	0.50	2.5
M. adsorption	1.85 ± 0.09 (1.85)	-5 ± 11 (3)	34 ± 4 (23)	0.61	2.9
2 + 2 CO	1.63 ± 0.05 (1.72)	97 ± 14 (109)	21 ± 5 (9)	0.43	3.5

^aVarying this parameter across all possible values did not result in an R -factor greater than the variance.

Table 3. Best fitting PhD structural models for single site occupation of a given DFT model for the PhD data taken after annealing the sample. The structural parameters of the PhD fit are given, the corresponding parameters from the DFT calculation are shown in brackets.

	R (Å)	ϕ (°)	θ (°)	R -factor	ΔF
EB-allyl	1.65 ± 0.05 (1.75)	57 ± 27 (66)	26 ± 4 (20)	0.40	2.6
DB-allyl	1.65 ± 0.04 (1.74)	82 ± 11 (83)	20 ± 4 (14)	0.44	2.4
EB-flagpole	1.69 ± 0.05 (1.71)	40 ± 10 (39)	31 ± 3 (26)	0.37	1.3
DB-flagpole	1.60 ± 0.04 (1.72)	8^a (10)	3 ± 8 (5)	0.33	2.5
M. adsorption	1.83 ± 0.16 (1.85)	3 ± 13 (3)	34 ± 5 (23)	0.52	2.8
2 + 2 CO	1.62 ± 0.04 (1.72)	114 ± 9 (109)	20 ± 3 (9)	0.31	3.3

^aVarying this parameter across all possible values did not result in an R -factor greater than the variance.

room temperature there remains a kinetic barrier to the formation of the flagpole structures that can be overcome by mild thermal annealing.

3.2. Qualitative analysis of PhD data

The oxygen 1s PhD data of acetophenone on Si(001) after room temperature deposition and after annealing the sample are shown in figures 2(a) and (b), respectively. Before performing a quantitative analysis it is useful to gain some insight from a qualitative inspection of the modulations. In particular, when the angle between the emitter, nearest-neighbour scatterer and detector is 180°, the modulation intensity will be at its greatest. Thus, if the emitter atom is adsorbed in a direct atop site (when an atom is directly above another, with no lateral displacement), the strongest modulations will be observed at normal emission. However, if the atom is off-atop, weaker modulations will be observed at normal emission, with stronger modulations at higher emission angles. Secondly, short emitter to nearest-neighbour scatterer distances result in long period modulations; long emitter to nearest-neighbour scatterer distances result in short period modulations. The O 1s modulations after room temperature deposition and after annealing (figure 2) are comparably weak ($\sim \pm 20\%$), and show the strongest modulations at an emission angle between 0° and 20° (with respect to normal emission). This suggests that the oxygen atoms are adsorbed in an off-atop site or sites. After annealing, the modulations still exhibit a relatively long period, again suggesting a comparably short oxygen-silicon bond length (< 2 Å). Inspection of the most energetically favourable DFT calculated

structures (figure 1) reveals that each of the computed structures has oxygen positioned in an off-atop site, in agreement with this qualitative inspection of the PhD data. Specifically, the oxygen atom in the DB-allyl is found 0.42 Å off-atop, EB-allyl 0.60 Å, DB-flagpole 0.16 Å, EB-flagpole 0.74 Å off-atop, molecular adsorption 0.71 Å off-atop and 2 + 2 CO 0.27 Å off-atop. Thus, the EB models adsorb with the oxygen atom more off-atop than the DB models, with EB-flagpole the furthest off-atop.

3.3. Quantitative PhD: multiple-scattering calculations

3.3.1. Room-temperature deposited. The best fits determined by multiple-scattering calculations of the six DFT structures as a single adsorption site are shown in table 2, with the corresponding values predicted by DFT shown in brackets. None of these models resulted in an R -factor less than 0.44, indicating poor agreement between the experimental and theoretical modulations in PhD. Furthermore, the agreement between the DFT and PhD structural parameters is poor for all models ($\Delta F > 2$) except for EB-flagpole ($\Delta F = 1.0$). This indicates that none of the structures in figure 1(a)–(f) on their own can account for the modulations observed in the PhD data. This is not particularly surprising, since previous STM work [16, 17] has already indicated that the surface should be at least a mixture of the two allyl features.

Therefore, we have also performed PhD modelling considering a surface composed of a various mixture of two out of the six configurations in figures 1(a)–(f). The results of these fittings can be found in the supplementary information (see table S1 in the ESI). Unfortunately, these two-site models resulted

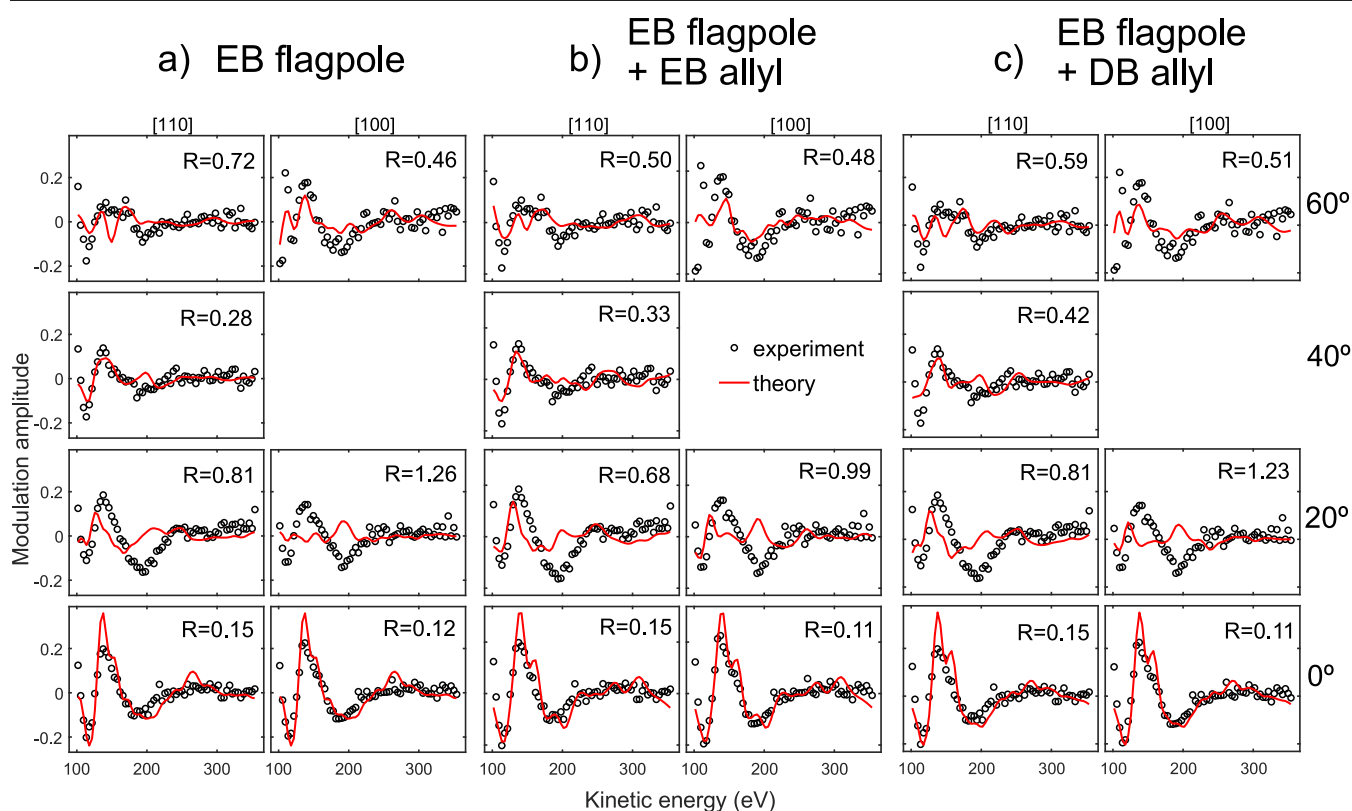


Figure 3. Comparison of the experimental and theoretical PhD modulations for the best fits found in the annealed preparation. Angles listed on the right-hand side are photoelectron emission angles with respect to normal emission. Inset is the individual R -factor for fit of each experimental PhD spectra to its corresponding theoretical spectra.

in only minor improvements in the R -factor (e.g. 0.43 for EB-allyl combined with EB-flagpole) and poor agreement of the structural parameters ($\Delta F > 2$). The best R -factor (0.39) is observed when EB-flagpole is combined with the 2 + 2 CO structure, but the agreement with DFT is poor ($\Delta F = 3$). The poor agreement, especially for the two allyl model, at first does seem surprising, since STM observations of the room temperature dosed surface show very clearly a surface composed primarily of two separate allyl features [16, 17]. However, resolution to this apparent discrepancy comes when the nature of the sample preparation is taken into account. In the STM experiments the surface is exposed to a very low dose of acetophenone in order to clearly identify isolated adsorbates. In contrast, in order to have a sufficient signal-to-noise ratio for the PhD experiments, the surface must be exposed to a higher dose of acetophenone. In performing our experiments, we observed the XPS signal from the dosed surface and we were careful to keep the dose sub-saturation; nevertheless, the higher coverage apparently results in sufficient percentage of additional configurations as to make accurate determination of the adsorbate structures by PhD fitting analysis prohibitive, and indeed prior studies have indicated that adsorption at high coverages results in the formation of multiple different local adsorption sites [18, 19].

Thus, we were not able to determine a good fit to confirm the adsorption structures of the room temperature dosed system; both single and two site models failed to adequately explain the experimental PhD data. Due to the ‘trial and error’

approach necessary for performing PhD data analysis, it is not realistic to model three or more differing structures, as adding in an ever greater number of fitting parameters is more likely to result in a greater number of non-unique solutions. Nevertheless, it is known that metastable structures, such as the 2 + 2 product, can be converted to the more stable structures by thermal anneal or STM activation [15], and as the average tilt angle of the molecule increases with thermal annealing, more space on the surface becomes available to convert these metastable structures into the more stable allyl or flagpoles structures. This suggests that mildly annealing the surface may increase the order and therefore also our chances of obtaining reliable fits to the PhD data.

3.3.2. Annealed. In good agreement with prior work, the single site allyl models result in both a poor R -factor (> 0.4) and $\Delta F (> 2)$ when used to fit to the annealed surface data. The best fits found for a single-site modelling of the DFT structures by multiple-scattering calculations are shown in table 3, and the corresponding predicted values by DFT for these parameters are shown in brackets. The best R -factors, 0.31 and 0.33, are found for the 2 + 2 CO and DB flagpole structures, respectively. However, the bond length in these structures (1.62 ± 0.04 Å and 1.60 ± 0.04 Å for 2 + 2 CO and DB-flagpole, respectively) is significantly shorter than that predicted by DFT (1.72 Å). Notably, the same models with a bond length of 1.72 Å have an R -factor of ~ 1.0 . As such, despite the good R -factor, neither the DB-flagpole nor the 2 + 2 CO structure

Table 4. Best fitting PhD structural models for two site occupation of a given DFT model for the PhD data taken after annealing the sample. The main structural parameters of the PhD fit are given, the corresponding parameters from the DFT calculation are shown in brackets.

	R (Å)	ϕ (°)	θ (°)	% occupation	R -factor	ΔF
EB-allyl + EB-flagpole	1.7 ± 0.2 (1.75)	70^a (66)	19^a (20)	49	0.35	1.9
	1.69 ± 0.04 (1.71)	40 ± 12 (39)	33 ± 3 (26)	51		
DB-allyl + EB-flagpole	1.72^a (1.74)	80^a (83)	10^a (14)	14	0.38	1.9
	1.69 ± 0.05 (1.71)	38 ± 12 (39)	32 ± 4 (26)	86		

^aVarying this parameter across all possible values did not result in an R -factor greater than the variance.

provide a good fit to our PhD data. The next best R -factor for our single site modelling is the EB-flagpole, with an R -factor of 0.37. In addition to an excellent R -factor, this configuration also has an excellent agreement between the PhD and DFT structural parameters ($\Delta F = 1.3$). The comparison between the experimental and theoretical PhD modulations is shown in figure 3(a).

Two-site modelling of the annealed system led to only one structure with a better R -factor than the single site EB-flagpole structure (see table S2). Specifically, a mixture of EB-allyl and EB-flagpole ($R_{\text{fac}} = 0.35$, $\Delta F = 1.9$). Only one other two site model resulted in a $\Delta F < 2$, which was a mixture of an DB-allyl and EB-flagpole ($R_{\text{fac}} = 0.38$). The comparison between the experimental and theoretical PhD modulations for both two site models are shown in figures 3(b) and (c) (respectively), and the structural parameters for the fits are shown in table 4.

4. Discussion

PhD is a powerful tool for the elucidation of atomic-scale structural parameters that are inaccessible to other techniques. However, it is most successful when applied to surfaces with a small number of distinct adsorption configurations. Thus, when applied to our room-temperature dosed surface, where multiple adsorption configurations exist, we were not able to obtain useful fits to our PhD data.

By annealing the surface, we were able to reduce the number of adsorbate structures, and fitting our PhD data to a two-species model produced an excellent fit to our data; in particular, the PhD data clearly indicates that the surface consists of a mixture of the EB-flagpole, and the EB-allyl adsorbate. Thus, our PhD data resolves the uncertainty that remained after our prior detailed XPS, NEXAFS, STM, and DFT studies [16–18], and identifies the EB-flagpole, and not the DB-flagpole, as the predominant structure formed by acetophenone on Si(001) after room temperature dosing and mild thermal annealing. It is worth noting that this is not the structure that might be anticipated by a naive interpretation of the DFT calculated structures, since the EB-flagpole is predicted to be less thermodynamically stable than the DB-flagpole. This can be attributed to the reaction pathway, from the allyl structures to the flagpole structures, being kinetically controlled.

It is important to note that, although it was possible to achieve a good fit to the combined data set, in none of the structures was a good fit to the 20° emission angle data found. In figure 3, the R -factors for each individual emission angle are shown, and universally the 20° emission angle has a notably

poor fit (>0.5). While the 40° and 60° emission angle data in some of the models also have a numerically poor fit, the comparatively small modulations ($\sim 10\%$) found at these emission angles will result in an underestimate in the quality of the fit at these emission angles. However the data at 20° are almost as strongly modulating as those at normal emission. This suggests that there is an aspect of the adsorbate structure that is not being well modelled within the structures that we have trialled. This could be indicative of a surface reconstruction, or the presence of a minority species on the surface that we have not modelled in our calculations. It is therefore important to qualify what this study was. Within this study we were explicitly comparing how well the theoretically predicted DFT structures match the experimentally measured structural data. Within this constraint, it is clear that the EB flagpole structure provides the best available fit to the experimental data, and thus likely dominates the surface post annealing.

The dominant structure upon room-temperature adsorption is the DB-allyl [16]. To convert the adsorbate from this allyl structure to the flagpole structure requires removing a H atom from the terminal C atoms of the acetyl group. There are two possible reaction pathways, one where the hydrogen abstraction occurs directly from the DB-allyl, and the other where the DB-allyl first converts to the EB-allyl [16]. Our results confirm that the latter is the kinetically preferred pathway. Indeed, direct manipulation of the DB-allyl using voltage pulses delivered by an STM tip has demonstrated the ability to induce the DB-allyl to convert to the EB-allyl, but the reverse process was never observed [16]. In addition, we note that the EB-allyl and EB-flagpole structures are found in DFT calculations to have the oxygen and carbon atoms of their acetyl groups in significantly more off-atop positions than the corresponding dimer-bridge structures, which is consistent with our observation of weak O 1s modulations. We speculate that the physical proximity of the terminal carbon of the acetyl group in relation to the neighbouring silicon atoms may result in a lowering of the activation energy for the H abstraction from the EB-allyl over the competing process of H abstraction from the DB-allyl, contributing to the preference for the reaction pathway resulting in the EB-flagpole structure.

5. Conclusions

We have presented an O 1s energy-scanned PhD study, with supporting DFT calculations, of the adsorption structures of acetophenone on the technologically pervasive silicon (001) surface. The unique capabilities of the PhD technique allow

us to answer a fundamental question regarding the physics and chemistry of this interaction that has remained elusive despite significant prior work with STM, DFT, XPS, and NEXAFS. In particular, we demonstrated that the most likely configuration adopted by the acetophenone adsorbate after room temperature dosing followed by a mild thermal anneal is an end-bridge flagpole configuration. This is despite DFT calculations suggesting that a 90 degree azimuthally-rotated structure, the dimer-bridge flagpole, is energetically favoured. Our results indicate that the surface reaction is kinetically-limited, and allow us to identify the preferred reaction pathway, where the acetophenone adsorbate transitions through the end-bridge allyl configuration. We hasten to add that this work has focussed on contrasting specific models predicted by DFT against our experimental PhD measurements, however both DFT and PhD are inherently imagination limited: if the basic structural parameters of the real structure are not present in our starting models, the real structure will not be found. With that limitation in mind, as interest in molecular adsorption on semiconductors progresses from small uni-functional adsorbates, to larger, technologically-relevant multi-functional adsorbates, we anticipate that PhD can become an important complementary tool for elucidation of structural properties inaccessible to other techniques and the development of molecular-functional devices.

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Data availability statement

The data that support the findings of this study are openly available via <https://zenodo.org> at [28].

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